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(54) Random copolymer and process for production thereof
Statistisches Copolymer und Verfahren zu dessen Herstellung
Copolymère statistique et son procédé de production

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(56) References cited:

EP-A- 0 156 464

US-A- 3 494 897

D. a.....

Description

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This invention relates to a novel random copolymer comprising at least polymerized units from ethylene and polymerized units from a cyclcolefin as monomers, and a process for production thereof. More specifically, it relates to a cyclcolefin type random copolymer having excellent transparency, thermal resistance, heat aging resistance, chemical resistance, solvent resistance, dielectric properties and mechanical properties and a narrow molecular weight distribution.

Polycarbonate, odiy(methyl methacrylate) and poly(ethylene terephthalate) are known as synthetic resins having excellent transparency. The polycarbonate resin has excellent heat resistance, heat aging resistance and impact strength as well as excellent transparency, but have the defect of poor chemical resistance since it is readily attacked by strong alkalis. Poly(methyl methacrylate) has the defect that it is susceptible to attack by ethyl acetate, acetone and tollene is swollen in ether, and has tow heat resistance. Polyethylene terephthalate has excellent heat resistance and mechanical properties but has the defect of possessing weak resistance to strong acids or alkalis and of being susceptible to hydrolysis.

Many polyolefins well known as general-purpose resins have excellent chemical resistance, solvent resistance and mechanical properties, but have poor neat resistance. Furthermore, they have poor transparency because they are crystalline. The transparency of polyolefins is generally improved by adding a nucleating agent to render the crystall structure fine, or by performing quenching to stop the growth of crystals, but such measures have not proved to be entirely effective. Father, the addition of a third component such as the nucleating agent is likely to impair the innerent excellent properties of the polyolefins. Furthermore, the quenching method requires large scale equipment, and is also likely to reduce heat resistance or rigidity with a decrease in crystallinity.

On the other hand, methods have been proposed for producing oclymers having superior transparency by coogreging ethylene with bulky componers. U.S. Patent 2,883,372 discloses a copplymer of ethylene with 2,3-dihydrodicyclopentaciene. This copplymer has a well balanced combination of rigidity and transparency but low heat resistance as shown by its glass transition temperature of about 160°C.

Jacanese Patent Publication No. 14,910/1971 discloses a process for producing a homo- or co-polymer having colymerized units represented by the following formula (a)

wherein R is hydrogen tha lower alkyligroup, which comprises colymerizing a 1.4.5.3-dimethano-1.2.3.4.4a.5.3.3a-i obtanydronachthalene represented by the following formula (b):

wherein R is as defined above

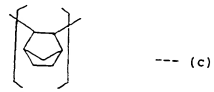
alone or with a cyclic diefin selected from styrene, adenaphthylene, bioyolo-[2,2,1]heptene-2, alkyl-substituted products of the heptene and cyclopentene in the presence of an alcohol as a reducing agent using a halide of a noble metal such as ruthenium as a catalyst.

Homo- or co-polymers comprising the 1,4,5,8-dimethano-1,2,3,4,4a,5,8 Ba-octanyoronabnthalene as part or the whole of monomers and methods for production thereof are also disclosed in Japanese Laid-Open Patent Publications 159,598/1975, 127,728/1983, 51,911/1984, 81,315/1984 and 81,316/1984 and U. S. Patent 4,178,424.

The colymers disclosed in these prior art documents are ring-opened polymers having polymerized units resulting from ring-opening of monomers as in the polymerized units represented by formula (a). As can be understood from formula (a), these ring-opened polymers are structurally characterized by the fact that the main polymer chain contains an unsaturated vinyl linkage. Because of the presence of the unsaturated vinyl linkage, these polymers have poor heat aging resistance.

All of the polymers described in the above prior an documents are ring-opened polymers obtained by the ring scission of a monomer. U. S. Patents 3,330,815 and 3,494,897 and Journal of Polymer Science: Polymer Physics Edition, volume 12, 1607-1618 (1974) describe polymers of the type obtained by the opening of the double bond of a monomer without the ring scission of the bicyclohept-2-ene skeleton of the monomer.

U. S. Patent 3,330,815 discloses that a polymer comprising polymerized units of the following formula (c)



or the substitution product thereof is obtained from bicyclo[2.2.1]hept-2-ene -2-ene of the following formula (d)

--- (a)

or its substitution product.

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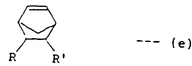
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U. S. Patent 3,494,897 discloses a process for producing alcopolymer of ethylene with a bicyclo[2,2,1]-nept-2-ene represented by the following formula (e)



Example 44 and claim 92 of this patent discloses a copolymer of ethylene with tetracyclo[6,2,1,1^{3,6},0^{2,7}]-4-dodecene of the following formula (f) encompassed within the formula (e)

--- (f)

and ethylene. The Patent does not describe the content of the tetracyclododecene of the copolymer of Example 42 When it is calculated under the assumption that all of the tetracyclododecene charged was introduced into the polymer the polymer contains polymerized units derived from about 2 mole% at most of the tetracyclododecene. U. S. Patent 3.494,897 does not at all describe specifically copolymers of ethylene with tetracyclododecenes other than the tetracyclof[5.2.1,13.6,02.7]-4-dodecene of formula (f).

Journal of Polymer Science: Polymer Physics Edition, volume 12, 1607-1618 (1974) discloses copolymers of ethylene and norbornene derivatives having dynamic mechanical properties. Table II of this document shows a copolymer of ethylene containing 1 mole% of polymerized units derived from the same compound as formula (f) (named octany-drodimethanonaphthalene in this document). Table IV thereof states that this copolymer has a beta-relaxation temperature of 15°C

The copolymers described in these prior art references which contain only 2% at the most of obtahydrodimethanonaphthalene have a glass transition temperature of about 15°C at the highest. They strongly exhibit a rubbery nature and have poor heat resistance and mechanical properties.

U.S. Patent Application Serial No. 695.151 (European Laid-Open Patent Application No. 0155454) discisses a random copolymer composed of ethylene, a 1.4.5.6-dimethano-1.2.3.4.4a.5.8.6a-octahydronaphthalene and option-the 1.4.5.8-dimethano-1.2.3.4.4a.5.8.8a-octahydronaphthalene and option-the 1.4.5.8-dimethano-1.2.3.4.4a.5.8.8a-octahydronaphthalene to polymerized units from ethylene is from 3.97 to 95 and the 1.4.5.8-dimethano-1.2.3.4.4a,5.8.6a-octahydronaphthalene is incorporated in the polymer chain as polymerized units without ring-opening.

The specification of U. S. Patent No. 2.883,372 discloses a normally solid copolymer of ethylene and 2.3-dinyd-rodicyclopentaciene having the following formula

said copolymer having a density in the range of 0.96 to 1.02 and being capable of being compression-molded at

The present invention seeks to provide a copolymer having excellent transparency, thermal resistance, heat aging resistance, chemical resistance, solvent resistance, dielectric properties and mechanical properties. Accordingly the present invention provides a random copolymer

which is transparent and is characterized in that

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(A) the copolymer comprises polymerized units from ethylene and polymerized units of at least one cycloolefin

 R^{5} R 9 $R^{\frac{2}{7}}$ (I) R¹⁰ 18 R 4 Ŕ

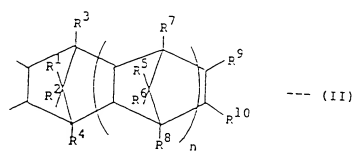
wherein $B^1, B^2, B^3, B^4, B^5, B^6, B^7$ and B^8 are identical or different and each represents a hydrogen atom, halogen atom or a hydrocarbon group having 1 to 10 carbon atoms, and R9 and R10 are identical or different and each represents a hydrogen atom, a halogen atom or a hydrocarbon group having 1 to 10 carbon atoms or \mathbb{R}^9 and \mathbb{R}^{10} are bonded to each other to form a group of the following formula

$$(R^{11} - C - R^{12})_{\ell}$$

in which R¹¹ and R¹² are identical or different and each represents a hydrogen atom, a halogen atom or a hydrocarbon group naving 1 to 10 carbon atoms, and ℓ is an integer of 3 or 4, and n is a positive integer of 1 to 3.

(B) the mole ratio of the polymerized cycloolefin units to the polymerized ethylene units is from 3:97 to 60:40.

(C) the or each cycloolefin of formula (i) is incorporated in the polymer chain as polymerized units represented by



wherein all symbols are as defined above.

(D) the copolymer has an intrinsic viscosity [η], measured in decalin at 135°C, of 0.01 to 20 dl/g.

(E) the copolymer has a molecular weight distribution (\overline{M} w/ \overline{M} n), measured by gel permeation chromatography, of

(F) the copolymer has a crystallinity, measured by X-ray diffractometry, of 0 to 10%, and

(G) the copolymer has a glass transition temperature (Tg) of 10 to 240°C as measured by a Dynamic Mechanical

55 The present invention also provides a process for producing a random copolymer as defined above, which process comprises continuously copolymerizing ethylene with the or each cycloolefin represented by formula (I) in a hydrocarbon medium in the presence of a catalyst formed from a vanadium compound soluble in the hydrocarbon medium and an organoaluminum compound while maintaining the atomic ratio of aluminum atoms to vanadium atoms in the reaction

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system, Al; V, at 2 or more: 1.
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The present invention additionally provides a process for preparing a modified random copolymer, which process comprises grafting on to a random copolymer, as defined above, 0.1 to 50 parts by weight of a monomer selected from (a) alpha.beta-unsaturated carboxylic acids and/or their derivatives, (b) styrenes, (c) organic silicon compounds having an olefinically unsaturated bond and a hydrolyzable group, and (d) unsaturated epoxy monomers per 100 parts by weight of the said random copolymer.

The present invention further provides molded articles prepared from a random copolymer as defined above or from a modified random copolymer prepared by a process as defined above.

The present invention also provides a modified random copolymer which obtainable by a process as defined above In formula (I), R1, R2, R3, R4, R5, R6, R7 and R5 are identical or different, and each represents a hydrogen atom, a halogen atom or a hydrocarbon group having 1 to 10 carbon atoms. The halogen atom is, for example, fluorine, chierine or bromine. The hydrocarbon group is preferably an alkyl group. Preferably the alkyl group has 1 to 5 carbon atoms. The alkyl group may be linear or branched, and includes, for example, methyl, ethyl, n-propyl, isopropyl, n-butyl, iso-butyl, sec-butyl, t-butyl, pentyl, hexyl, heptyl, octyl, nonyl and decyl.

Examples of the halogen atom and the hydrocarbon group represented by ${\sf R}^9$ and ${\sf R}^{10}$ are the same as those given above.

The integer n is preferably 1 or 2. Examples of the cycloplefin of formula (I) include

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             1.4,5.5-dimethano-1,2.3,4.4a,5.5.8a-cctahydronaphthalene,
            2-methyl-1,4,5,5-dimethano-1,2,3,4,4a,5,8,8a-octahydronaphthalene.
            \hbox{2-ethyl-} \hbox{1.4,5,5-dimethano-} \hbox{1.2,3.4,4a.5.8.8a-octahydronaphthalene},
            2-propyi-1.4.5.8-dimethano-1.2.3,4.4a.5.8.8a-octahydronaphthalene,
            2-hexyl-1,4,5,8-cimetnano-1,2,3,4,4a,5,8,8a-octahydronaphthalene,
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            2-stearyl-1,4,5,8-dimethano-1,2,3,4,4a,5,8,8a-octahydronaphthalene,
            2.3\text{-}dimethyl-1.4,5.8\text{-}dimethano-1,2.3,4,4a,-5.8.8a-octahydronaphthalene.}\\
            2-methyl-3-ethyl-1,4,5,8-dimethano-1,2,3,4,4a,5,8,8a-octahydronaphthalene,
            2-chloro-1,4,5,8-dimethano-1,2,3,4,4a,5,8,8a-octahydronaphthalene,
            2-bromo-1,4,5,3-dimethano-1,2,3,4,4a,5,8,8a-octahydronaphthalene,
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            2-fluoro-1,4,5,8-dimethano-1,2,3,4,4a,5,8,6a-octahydronaphthalene,
            2.3-dichicro-1,4,5,8-dimethano-1,2,3,4,4a,-5,8,8a-octahydronaphthalene,
            2-cyclohexyl-1,3,5,5,-dimethano-1,2,3,4,4a,-5,8,8a-octanydronaphthalene,
            2-n-butyl-1,4,5,8-dimethano-1,2,3,4,4a,5,8,6a-octahydronaphthalene,
            2-isobutyi-1,4,5,8-dimethano-1,2,3,4,4a,5,8,6a-octahydronaphthalene,
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            5.10-dimethyltetracyclo[4,4,0,12.5,17.10]-3-dodecene,
            2.10-dimethyltetracyclo[4,4,0,1<sup>2,5</sup>,1<sup>7,10</sup>] -3-dodecene,
            11,12-diwethyltetracyc1c[4,4,0,12.5,17.10]-3-dodecene.
           2.7.9-trimetnyltetracyclo[4.4.0,13.5,17.10]-3-dodecene.
           9-etnyl-2.7-aimetnyltetracyclo[4,4,0,1<sup>2,5</sup>, 1<sup>7,10</sup>]-3-dodecene,
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           9-isoputyl-2,7-dimethyltetracyclo-[4,4,0,12.5,19.10]-3-dodecene.
           9.11.12-trimethyltetracyclo[4,4,0,12.5,17.10]-3-dodecene
           9-ethyl-11, 12-dimethyltetracyclo[4,4,0,12.5, 17,10]-3-dodecene,
           9-isobutyl-11,12-dimethyl-3-tetracyclo-[4,4,0,12.5,17.10]-3-dodecene.
           5.3,9.10-tetramethyltetracyc1o(4,4,0,12.5, 17.10) -3-dodecene.
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           hexacyclo[5,5,1,13,6,110,13,02,7,09,14]-4-heptagecene.
           12-methylnexacyclo[6,6.1,13.6,110.13,02.7,09.14]-4-heptadecene.
           12-ethylhexacyclo[5,6,13,6,110,13,03,7,05,14]-4-neptadecene.
           12-isobutylhexacycio[6,6,1,13,6,110,13,C27, 09,14]-4-heptadecene
           1,5,10-trimethyl-12-isobutylnexacyclo[6,5,1, 13,5,110,13,02,7,09;14]-4-heptadecene.
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           cctacyclo[8.8,12.9,14.7,111.19,113.16,0,03.8, 012.17]-5-dococene,
           15-methylociacyclo(8.5 12.9 14.7 111.15 113.16 0.03.6 012.17)-5-coccene,
           15-ethyloctacycic(8.8.12.5.14.7.111.18.113.16.113.16.0.03.8.012.17]-5-dococene.
           1.3-dimethylpentacyclo[6.6.1,13.5,02.7,09.14]-4-hexadecene,
           1.6-aimethylpentacyclo[5.6.1,1<sup>3.6</sup>,0<sup>2.7</sup>,0<sup>9.14</sup>]-4-hexadecene.
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           15.16-dimethylpentacyclo[6.6.1,13.6]02.7]09.14]-4-hexadecene.
           pentacyclo[6.5,1,13.6,02.7,09.13]-4-pentadecene.
           1,3-dimethylpentacyclo[6.5,1,1<sup>3,6</sup>,0<sup>2,7</sup>,0<sup>9,13</sup>]-4-pentadecene
           1.5-dimetry/pentacyclo[6.5.1,13.6,02.7,09.13]-4-pentadecene.
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14,15-dimethylpentacyclo[6,5,1,1^{3,6},0^{2,7},0^{9,13}]-4-pentadecene. pentacyclo[6,6,1,13,6,02,7,09,14]-4-hexadecene, heptacyclo[8.7, 12.9, 14.7, 111.17, 0, 03.8, 012.16]-5-eicosene, and pentacyclo[8.8, 12.9, 14.7, 111.18,0,03.8,012.17]-5-heneicosene.

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The compounds of formula (I) may be used singly or in combination.

The compounds of formula (I) can be easily produced by condensing cyclopentadienes with the corresponding olelins by the Diels-Alder reaction.

According to the process of this invention, the cycloolefin of formula (I) is copolymerized continuously with ethylene in a hydrocarbon medium in the presence of a catalyst. The catalyst is one formed from a soluble vanadium compound and an organization compound. In the copolymerization reaction carried out continuously, the atomic ratio of alu-

The soluble variadium compound used in the process of this invention as a component of the catalyst is a variadium. compound soluble in the hydrocarbon medium in the polymerization reaction system. It may include variadium compounds of formula $VO(OR)_a X_b$ or $V(OR)_c X_d$ wherein R is a hydrocarbon group such as an aliphatic, alicyclic or aromatic hydrocarbon group having 1 to 20 carbon atoms, preferably an aliphatic hydrocarbon group having 1 to 5 carbon atoms. especially 1 to 3 carbon atoms, 0≦a≦3, 0≦b≦3, 2≦a+b≦3, 0≦c≦4, 0≦d≦4, 3≦c+d≦4, and X is halogen such as Cl. Br or I and adducts of these compounds with electron donors. Specific examples are VOCl₂, VO(OCH₃)Cl₂, VC $(\text{OCH}_3)_5, \ \ \text{VO}(\text{OC}_2\text{H}_5)_{1.5}\text{Cl}_{1.5}, \ \ \text{VO}(\text{OC}_2\text{H}_5)\text{Cl}_2, \ \ \text{VO}(\text{OC}_2\text{H}_5)_2\text{Cl}, \ \ \text{VO}(\text{O-iso-C}_3\text{H}_7)\text{Cl}_2, \ \ \text{VO}(\text{O-iso-C}_$ $C_{3}H_{7})_{1.5}Cl_{1.5}, VO(O-n-C_{3}H_{7})_{1.5}Cl_{1.5}, VO(O-n-C_{3}H_{7})Cl_{2}, VO(O-n-C_{2}H_{7})Cl, VO(O-C_{5}H_{11})_{1.5}Cl_{1.5}, VO(O-n-C_{4}H_{9})Cl_{2}, VO(O-n-C_{5}H_{11})_{1.5}Cl_{1.5}, VO(O-n-C_{5}H_{11})_{1.5}Cl_{1.5}, VO(O-n-C_{5}H_{11})_{1.5}Cl_{1.5}, VO(O-n-C_{5}H_{5})Cl_{5}, VO($ (O-n-C₄H₉)₂Cl, VO(O-iso-C₄H₉)₂Cl, VO(O-sec-C₄H₉)₃. VO(OC₂H₅)₃. VOBr₂. VCl₄. VOCl₂, VCl₃. VBr₄. VBr₅. VO(O-n-

Compounds having at least one AI-C-bond in the molecule can be used as the organoaluminum compound used as a catalyst component. Examples are

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(i) organizationinum compounds of formula

wherein E¹³ and E¹⁴ are identical or different and each represents a hydrocarbon group having 1 to 15 carbon atoms, preferably 1 to 4 carbon atoms. X represents a halogen atom such as CI, Brior I, mis a number represented by 0≦m≦3, t is a number represented by 0≦t<3, p is a number represented by 0≦t<3, and q is a number repre-

(ii) Complex alkylated compounds of metals of Group I of the periodic table and aluminum of formula

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wherein M^{ϵ} is Li. Na or K, and B^{15} is as defined above

Examples of the organical uninum compounds (i) are compounds of formula

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wherein R^{13} and R^{14} are as defined above and m is preferably a number represented by 1.5≦m≦3, compounds of formula

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wherein \mathbb{R}^{13} is as defined above. X is halogen such as Cl. Br or I, and m is preferably 0<m<3, compounds of formula

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wherein R¹³ is as defined above and m is preferably a number represented by 2≲m<3, and compounds of formula

wherein R¹³ and R¹⁴ are as defined above. X is halogen such as CI. Br or I, 0<m≤3, and 0≤q<3, provided that

Examples of the aluminum compounds (i) include trialkyl aluminums such as triethyl aluminum, triisopropyl aluminum and tributyl aluminum; dialkyl aluminum alkoxides such as diethyl aluminum ethoxide and dibutyl aluminum butoxide; alkyl aluminum sesquialkoxides such as ethyl aluminum sesquiathoxide and butyl aluminum sesquibutoxide partially alkoxylated alkyl aluminums having the average composition of $R_{0.5}^1 A_1(\text{OR}^2)_{0.5}$; partially halogenated alkyl aluminums, for example dialkyl aluminum halides such as diethyl aluminum chloride, dibutyl aluminum chloride and ethyl aluminum bromide, alkyl aluminum sesquihalides such as ethyl aluminum sesquichloride, butyl aluminum sesquichloride and ethyl aluminum sesquibromide, and alkyl aluminum dihalides such as ethyl aluminum dichloride, procylatininum cichioride and butyl aluminum dibromide; partially hydrogenated alkyl aluminums, for example dialkyl aluminum hydrides such as diethyl aluminum hydride and dibutyl aluminum hydride, and alkyl aluminum dihydrides such as ethyl aluminum dihydride and propyl aluminum dihydride; and partially alkoxylated and halogenated alkyl aluminums such as ethyl aluminum ethoxychloride, butyl aluminum butoxychloride and ethyl aluminum aluminum ethoxybromide. As compounds similar to (i), organoaluminum compounds having at least two aluminums bonded through an oxygen or nitrogen atom may also be used. Examples of such compounds are $(C_2H_5)_2AlOAl(C_2H_5)_2$, $(C_4H_9)_2AlOAl(C_4H_9)_2$ and $(C_2H_5)_2AlOAl(C_2H_5)_2$.

Examples of the compounds (ii) are $LiAl(C_2H_5)_4$ and $LiAl(C_7H_{15})_4$.

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Among these, the aikyl aluminum halides, alkyl aluminum dihalides and mixtures of these are especially preferred. The copolymerization reaction in accordance with the process of this invention is carried out in a hydrocarbon medium. Examples of the solvent include aliphatic hydrocarbons having 5 to 15 carbon atoms such as pentane, hexame neptane, octane and kerosene; alicyclic hydrocarbons having 5 to 15 carbon atoms such as cyclopentane, cyclohexame and methylcyclonexane; and aromatic hydrocarbons having 6 to 15 carbon atoms such as benzene, toluene and xylene. These solvents may be used singly or in combination.

in the process of this invention, the copolymerization reaction is carried out continuously. During the reaction, the ratio of aluminum atoms to variadium atoms, AltV, is maintained at 2 or more:1, preferably 2 to 50:1, especially preferably 3 to 20:1.

The soluble variadium compound and the organizationium compound are fed to the reaction system usually after dilution with the hydrocarbon medium. The concentration of the vanadium compound to be fed to the polymerization reaction system is usually not more than 10 times, preferably 7 to 1 times, more preferably 5 to 1 times, that in the polymerization reaction system. On the other hand, the organization compound may be fed to the polymerization system after its concentration is adjusted to not more than 50 times that in the polymerization system. In the process of this invention, the concentration of the soluble variadium compound in the copolymerization reaction system is usually 0.01 to 5 gram-atoms/liter, preferably 0.05 to 3 gram-atoms/liter, as vanadium atoms, in the process of this invention for producing the cycloolefin random copolymer, the-copolymerization reaction may be carried out in the copresence of an electron donor in the polymerization reaction system in addition to the soluble vanadium compound and the organisaluminum compound as catalyst ingredients. Examples of the electron donor include oxygen-containing electron conors such as alcohols, phenols, ketones, aldehydes, carboxylic acids, esters of organic acids or inorganic acids, ethers, acid amides, acid annydrides and alkoxysilanes, and nitrogen-containing electron donors such as ammonia, amines, nitriles and isocyanates. The copolymerization reaction is carried out usually at -50 to 100°C, preferably -30 to 50°C more preferably -20 to 60°C. The average residence time in the copolymerization reaction differs depending: upon the type of the polymerization material, the concentrations of the catalyst components, and the polymerization temperature. Usually it is 5 minutes to 5 hours, preferably 10 minutes to 3 hours. The pressure during the copolymerization reaction is usually more than 0 to 50 kg/cm², preferably more than 0 to 20 kg/cm². If desired, it is carried out in the presence of an inertigas such as nitrogen or argon. To control the molecular weight of the copolymer, a molecular weight controlling agent such as hydrogen may be present in the reaction system. The polymerization reaction mixture after the copolymerization is treated in accordance with conventional methods to give the desired cycloolefin random copolymer

The mole ratio of ethylene cycloolefin fed to the copolymerization reaction system in the process of this invention is usually from 99:1 to 1:99, preferably 98:2 to 2:98.

in this invention, the cycloolefin is copolymerized with ethylene. As required, another copolymerized unsaturated monomer within a range which does not impair the objects of the invention, for example in an amount of 50 mole% preferably 40 mole%, based on the ethylene units in the copolymer may be copolymerized in addition to these two essential components. Specific examples of the other copolymerizable unsaturated monomer include alpha-olefins having 3 to 20 carbon atoms such as propylene, 1-butene, 4-methyl-1-pentene, 1-hexane, 1-octene, 1-decene, 1-decene, 1-tetradecene, 1-hexadecene, 1-cotadecene and 1-eicosane which are to be used in an amount of less than 1 mole per mole of the ethylene units in the resulting random copolymer; and cycloolefins such as cyclopentene, cyclohexane, 3-methylcyclohexane, cyclooctene, 3a.5.5.7a-tetrahydro-4,7-methano-1H-indene, nonconjugated dienes such as 1,4-hexadiene, 4-methyl-1,4-hexadiene, 5-methyl-1,4-hexadiene, 1,7-octadiene, dicyclopentadiene, 5-ethylidene-2-norbornene and 5-vinyl-2-norbornene, and norbornenes such as norbornene-2, 5-methylnorbornene-2, 5-i-butylnorbornene-2, 5-i-butylnorbornene-2, 5-6-dimethylnorbornene-2, 5-chloronorbornene-2, 2-fluoronorbornene-2, and 5.6-dichloronorbornene-2

The copolymerization reaction in accordance with this invention can be advantageously carried out by continuously

feeding ethylene, the cycloolefin, and optionally the other copolymerizable component as materials to be polymerized the soluble vanadium compound and organoaluminum compound as catalyst components, and the hydrocarbon medium into the polymerization system, and continuously withdrawing the polymerization reaction mixture from the polymerization reaction system.

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According to the continuous process of this invention, a random copolymer having a narrow molecular weight distribution and good transparency can be produced.

The resulting copolymer solution continuously withdrawn from the polymerization reaction system is a hydrocarbon medium solution of the random copolymer. The concentration of the random copolymer in the resulting copolymer solution is usually 2.0 to 20.0% by weight, preferably 2.0 to 10.0% by weight. Contacting the resulting copolymer solution with a ketone or alcohol gives a precipitate of the random copolymer. The precipitate is separated by such secarating means as filtration or centrifugal separation to give the random copolymer of this invention. The proportion of the ketone or alcohol used is usually 200 to 1,000 parts by weight, preferably 300 to 500 parts by weight, per 100 parts by weight of the resulting copolymer solution. Examples of the ketones are those having 3 to 7 carbon atoms such as acetone, methyl etnyl ketone, dietnyl ketone, methyl isobutyl ketone, dipropyl ketone and acetylacetone. Examples of the alcohols are alcohols having 1 to 4 carbon atoms such as methanol, ethanol, propanol, isopropanol butanol, sec-butanol and tert-butanol. The ketones or alcohols may contain a small amount of water. The use of a mixture of the ketone or alcohol with usually 0.1 to 10% by weight, preferably 1 to 5% by weight, of water is suitable since it gives a cooolymer powder having low contents of the unreacted cyclcolefin monomer and a low-molecularweight polymer and high bulk density. Contacting of the resulting copolymer solution with the ketone or alcohol is carried out with stirring at a temperature of usually 0 to 100°C, preferably 10 to 70°C, especially preferably at a temperature near the coiling point of the ketone or alcohol. This is effected by using a vessel-type mixer having agitating vanes such as turbine blades and comb blades. The rotating speed of the blades is usually 200 to 2,000 rpm, preferable, 800 to 1,500 rpm

As a result of the after-treatment described above, the random copolymer is precipitated usually in powder form to form a suspension. The suspension containing the copolymer is separated into the copolymer and the mother liquod such a separating means as centrifugation or filtration. The separated copolymer is further extracted with a ketone to give a random occolymer having low contents of the unreacted cycloolefin monomer and a low-molecular-weight distribution, a narrow composition distribution, and a high glass transition temperature. The proportion of the ketone used in the extraction is such that the concentration of the copolymer in the ketone is usually 10 to 100 g/liter, preferably 30 to 60 g/liter. The temperature at which the copolymer is contacted with the ketone is usually 10 to 55°C, preferably 15 to 50°C. The contacting treatment between the copolymer and the ketone is usually carried out with stirring using, for example, a vessel-type mixer having agitating blades such as turbine blades and come ciades. The rotating speed of the blades at this time is usually 50 to 400 rpm, preferably 100 to 200 rpm. The time required for contacting is usually 1 to 8 hours, preferably 2 to 5 hours. Examples of the ketone may be the same as those exemptified for the ketone used for the precipitation treatment. The copolymer which has been contacted with the ketone may be separated by separating means such as centrifugation or filtration. As required, the separated copolymer is further washed with the ketone. The washing conditions may be the same as the ketone contacting treatment conditions described above.

The random occorymer of this invention obtained by the above after treatment usually has a bulk density of 0.05 to 0.3, preferably 0.1 to 0.25, and is easy to handle at the time of molding. Furthermore, since the copolymer of this invention has low contents of the unreacted cycloolefin monomer and a low-molecular-weight polymer, optical materials, padicularly octical memory discs, molded from it permit reduction of noise caused by, for example, reflection

Thus, the present invention gives a random copolymer in which the mole ratio of polymerized cycloolefin of formula (i) units to polymerized ethylene units is from 3:97 to 60:40, preferably from 5:95 to 50:50

The random opposymer of this invention has an intrinsic viscosity [η], measured in decalin at 135°C, of 0.01 to 20 d/g preferably 0.05 to 10 d/g.

The random copolymer of this invention has a molecular weight distribution $(\overline{M} w | \overline{M} n)$, measured by gel permeation distribution ($\overline{M} v | \overline{M} n$), measured by gel permeation distribution ($\overline{M} v | \overline{M} n$), of not more than 4, preferably not more than 3. The random copolymer of this invention is characterized in that this molecular weight distribution is narrow. This means that the random copolymer of this invention has a low content of a low-molecular-weight polymer. Accordingly, when the random copolymer of this invention is molded into an optical material such as an optical memory disc or optical fibers, the optical material has good surface smoothness, low surface tack, or good mechanical properties and is suitable for optical applications.

The random copolymer of this invention has a crystallinity, measured by X-ray diffractometry, of 0 to 10%, preferably 0 to 5%. especially preferably 0 to 7%. The advantage of the copolymer having a crystallinity of 10% or below is that when it is moided into an optical material such as an optical memory disc or optical fibers, noise of the optical material caused by for example, refraction or diffused reflection is reduced.

The copplymer has a Tg of 10 to 240°C, more preferably 20 to 200°C

The random copolymers of this invention are molded by known mathods. For example, they can be extrusion-molded, injection-molded, blow-molded, or rotationally molded by using, for example, a veni-type extruder, a twin-screw extruder, a Cokneader, a plsticator, a mixtruder, a twin conical screw extruder, a gear-type extruder or a screwless extruder. In the molding process, known additives such as heat stabilizers, light stabilizers, antistatic agents, slip agents, antiblocking agents, antinaze agents, lubricants inorganic and organic fillers, dyes and pigments may be used as required.

Phenolic or sulfur-type antioxidants may be cited as examples of such additives. The phenolic antioxidants include for example, phenolis such as 2.6-di-ten-butyl-p-cresol, stearyl (3.3,-dimethyl-4-hydroxybenzyl)thiogiyoolate, stearyl beta-(4-hydroxy-3.5-di-ten-butyl-4-hydroxybenzyl)phosphate, 2.4,6-tris(3;5'-di-ten-butyl-4'-hydroxybenzyl)phosphate, 2.4,6-tris(3;5'-di-ten-butyl-4'-hydroxybenzyl)maionate, 2.2'-methyl-en-bis(4-methyl-6-ten-butylphenol), 4,4'-methylenebis(2.6-di-ten-butylphenol), 2,2'-methylenebis[5-(1-methylcyclo-hexyl)p-cresol], bis[3.5-bis[4-hydroxy-3-ten-butylphenyl)butyric acid] glycol ester, 4,4'-butylidenebis[6-ten-butyl-methyl-6-(2-hydroxy-3-ten-butylphenyl)butene, bis[2-ten-butyl-4-methyl-6-(2-hydroxy-3-ten-butyl-5-methylbenzyl)phenzyl] terephthalate, 1,3.5-tris(2.6-di-methyl-3-hydroxy-4-ten-butyl-benzyl) isocyanurate, 1,3.5-tris(3.5-di-ten-butyl-4-hydroxybenzyl) isocyanurate, 1,3.5-tris(3.

Examples of the sulfur-type antioxidant include dialkyl thiodipropionates such as dilauryl, dimyristyl or disteany thiodipropionate, and esters (such as pentaerythritol tetralaurylthiopropionate) formed between alkylthiopropionic acids such as butyl-, octyl-, lauryl- or stearylthiopropionic acid and polyhydric alcohols (such as glycerol, trimethylolethane trimethylolpropane, pentaerythritol or tris-hydroxyethyl diisocyanurate).

Phosphorous-containing compounds may also be incorporated. Examples include tricityl phosphite, trilauryl phosphite tridecyl phosphite, octyldiphenyl phosphite, tris(2,4-di-ten-butylphenyl) phosphite, tripnenyl phosphite, tris(0,4-di-ten-butylphenyl) phosphite, tris(0,4-di-ten-butylphenyl) phosphite, tris(0,4-di-ten-butylphenyl) phosphite, tris(0,4-di-ten-butyl-4-hydroxy-pnenyl) phosphite, tetra(C₁₂-C₁₅ mixed alkyl)-4,4'-isopropylidenediphenyl diphosphite tetrartridecyl)-4,4'-butylidenedis(3-methyl-5-tent-butylphenol) diphosphite, tris(3,5-di-tent-butyl-4-hydroxypnenyl, phosphite, tris(mono-di-mixed nonylphenyl) phosphite, hydrogenated 4,4'-isopropylidene diphonol pclyphosphite, discordiphonol), bis(4,4'-butylidenebis(3-methyl-6-tent-butylphenol)), 1,6-hexanediol diphosphite, phenyl-4,4'-isopropylidenedionenol-pentaerythritol diphosphite, bis(2,4-di-ten-butylphenyl)pentaerythritol diphosphite, bis(2,6-di-tent-butylphenyl)pentaerythritol diphosphite, tris(4,4'-isopropylidenebis(2-tent-butylphenol)), phosphite, phenyldiscordiphosphite, di(nonylphenyl)pentaerythritol diphosphite, tris(1,3-di-stearoyloxyisopropyl), phosphite, 4,4'-isopropylidenebis(2-tent-butylphenol)-di(nonylphenyl), phosphite, 9,10-dihydro-9-oxa-10-phosphaphenanthrene-10-oxide and letrakis(2,4-di-tent-butylphenyl)-4,4'-biphenylene diphosphonite.

There can also be used 6-hydroxycoumarone derivatives such as alpha-, beta-, gamma- and delta-tocopherois and mixtures thereof, a 2.5-dimethyl-substitution product, 2.5.8-trimethyl-substituted product or 2.5.7.8-tetramethyl-substituted product of 2-(4-methyl-pent-3-eny)-6-hydroxycoumarone, 2.2.7-trimethyl-5-tert-butyl-6-hydroxycoumarone, 2.2.5-trimethyl-6-tert-butyl-6-hydroxycoumarone, and 2.2-dimethyl-5-tert-butyl-6-nydroxycoumarone

It is also possible to incorporate a compound represented by the general formula

$$M_x Al_y (OH)_{2x+3y-2z} (A)_z .aH_2 O$$

wherein M represents Mg. Ca or Zn, A is an anion other than a hydroxyl anion, x, y and z are positive numbers, and a represents 0 or a positive number

Examples of the compounds of the above formula are

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 $Zn_6Al_2(OH)_{16}CO_3 \ 4H_2O.$ $Zn_6Al_2(OH)_{16}SO_4 \ 4H_2O.$ $Mg_6Al_2(OH)_{16}SO_4 \ 4H_2O.$

and

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Mg₆Al₂(OH)₁₂CO₃ 3H₂O

There may also be added an antioxidant having a 2-benzofuranone skeleton, such as 3-phenyl-2-benzofuranone and 3-phenyl-4,6-di-t-butyl-2-benzofuranone as disclosed in the specification of Japanese Laid-Open Patent Publication No. 501.181/1980

Examples of the light stabilizers include hydroxybenzophenones such as 2-hydroxy-4-methoxybenzophenone. 2-hydroxy-4-n-octoxybenzophenone, 2,2'-dihydroxy-4-methoxybenzophenone and 2,4-dihydroxybenzophenone; benzotriazoles such as 2-(2'-hydroxy-3'-fert-butyl-5'-methylphenyl)-5-chlorobenzotriazole, 2-(2'-hydroxy-3'-5'-di-tert-butyl-phenyl)-5-chlorobenzotriazole, 2-(2'-hydroxy-3'-5'-di-tert-butyl-nyl)benzotriazole; benzoates such as phenyl salicylate, p-tert-butylphenyl salicylate, 2,4-di-tert-butylphenyl-3,5-di-tert-butyl-4-hydroxybenzoate; nickel compounds such as 2,2'-thiobis (4-tert-octylphenol) nickel salt. [2,2'-thiobis(4-tert-octylphenolate)]-n-butylamine nickel salt and (3,5-di-tert-butyl-4-ny-croxybenzyl)-phosphonic acid monoethyl ester nickel salt; substituted acryloniriles such as methyl alpha-cyano-beta-oxalic acid diamide and N-2-ethylphenyl-N'-2-ethoxyphenyl oxalic acid diamide; and hindered amine compounds such as bis(2,2,6,6-tetramethyl-4-piperidine)sebacate, poly[(6-(1,1,3,3-tetramethylbutyl)imino)-1,3,5-triazine-2,4-diyl-4-ny-droxy-2,2,6,6-tetramethyl-1-piperidyl)ethanol.

Examples of the lubricants include aliphatic hydrocarbons such as paraffin wax, polyethylene wax and polypropylene wax; higher fatty acids such as capric acid, lauric acid, myristic acid, palmitic acid, margaric acid, stearic acid, acid acid acid acid behenic acid; metal salts of higher fatty acids such as lithium, calcium, sodium, magnesium and potassium salts of the above-exemplified fatty acids; aliphatic alcoholes such as palmityl alcohol, cetyl alcohol and stearyl alcohol; aliphatic amides such as caproic amide, caprylic amide, carpric amide, fluorine amide, myristic amide, calmitic amide and stearic amide; esters formed between fatty acids and alcohols; and fluorine compounds such as fluoroalkylcarcoxylic acids metal salts thereof, and metal salts of fluoroalkyl-sulfonic acids.

Examples of the fillers include inorganic or organic fibrous fillers such as glass fibres, silver- or aluminum-coated glass fibers, stainless steel fibers, aluminum fibers, potassium titanate fibers, carbon fibers, whiskers, Keviar® fibers and supernign elastic polyethylene fibers; and inorganic or organic powdery, granular or flaky fillers such as talc, calcium carbonate, magnesium nydroxide, calcium oxide, magnesium sulfate, graphite, nickel powder, silver powder, copper powder, carbon black, silver-coated glass beads, aluminum-coated glass beads, aluminum flakes, stainless steel flakes

The random copolymers of this invention have excellent transparency, heat resistance, heat aging resistance, chemical resistance, solvent resistance, dielectric properties and mechanical properties, a narrow and uniform molectural weight distribution, and a narrow and uniform composition distribution. Accordingly, those having a low molecular weight are synthetic waxes and are useful as candles, impregnating agents for matches, paper finishing agents, sizes, antioxicants for rubber, waterproofing agents for cardboards, slow releasing agents for chemical fertilizers, heat accumulators, binders for ceramics, paper condensers, electric insulating materials for electric wires and cables, agents for decelerating neutrons, fiber finishing aids, water-repellents for building materials, protecting agents for coatings, colishing agents, thixotropy imparting agents, core hardening agents for pencils and crayons, carbon ink substrates electrophotographic toners, lubricants and releasing agents for molding of synthetic resins, resin coloring agents, hotelessives, and lubricant greases. Those having a high molecular weight can be used in an optical field as optical lenses, optical discs, optical fibers and windowpanes, in an electrical field as a water tank for electrical irons, parts of electronic ovens, base boards for liquid crystal display, base boards for printed circuit boards, high frequency circuit boards and transparent electrically conductive sneets or films, in medical and chemical fields as syringes, pipettes and animal gages, and in other various fields as housings of measuring instruments, and helmets

When the copolymers of this invention have a glass transition temperature (Tg) below 100°C, they can be used in a field utilizing their shape memory, and as a vibration controlling material or a tube. Specifically, they can be used as joints for irregularly-shaped pipes, laminating agents for the interior and exterior parts of pipes and rods, clamping pins for optical fiber connectors, plaster casts, containers, automobile bumpers, various space eliminating materials, vibration controlling materials or sound insulating materials in the form of a laminate with a metal surface material, and tubes for use in medical therapy.

EP 0 203 799 B1

The random copolymers of this invention may be used as a blend with various known polymers. Examples of such known polymers are:

(A) Polymers derived from hydrocarbons having 1 or 2 unsaturated bonds

Polyolefins, such as polyethylene, polypropylene, polyisobutylene, poly(methylbutene-1), poly(4-methylpentene-1), poly(methylbutene-1), polyisoprene, polybutadiene and polystyrene, or crosslinked products thereof

Copolymers of the monomers constituting the above polymers with each other, such as ethylene/propylene copolymer, propylene/butene-1 copolymer, propylene/isobutylene copolymer, styrene/isobutylene copolymer, styrene copolymer rene/butadiene copolymer, terpolymers of ethylene, propylene and dienes such as hexadiene, bicyclopentadiene and 5-ethylidene-2-norbornene, and terpolymers of ethylene, butene-1 and dienes such as hexadiene, dicyclopen-

Blends, grafted polymers, and block copolymers of these polymers may also be cited.

(B) Halogen-containing vinyl polymers

Polyvinyl chloride, polyvinylidene chloride, polyvinyl fluoride, polychloroprene, and chlorinated rubbers.

(C) Polymers derived from alpha beta-unsaturated carboxylic acids or the derivatives thereof Polyacrylate, polymethacrylate, polyacrylamide, and polyacrylonitrile.

Copolymers of the monomers constituting the above-exemplified polymers with other copolymerizable monomers, such as acrylonitrile/butadiene/styrene copolymer, acrylonitrile/styrene copolymer, and acrylonitrile/sty-20

(D) Polymers derived from unsaturated alcohols, amines, acyl derivatives thereof, or acetals

Polyvinyl alcohol, polyvinyl acetate, polyvinyl stearate, polyvinyl benzcate, polyvinyl maleate, polyvinyl butyra: polyallyi phthalate and polyallyimelamine.

Copolymers of the monomers constituting the above-exemplified polymers with other copolymerizable monomers, such as ethylene/vinyl acetate copolymer.

(E) Polymers derived from epoxides

Polyethylene oxide and polymer derived from bisglycidyl ether.

(F) Polyacetais

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Polyoxymethylene, polyoxyethylene, and polyoxymethylene containing ethylene oxide.

- 35 (G) Polyphenylene oxide
 - (H) polycarbonate
 - (i) Polysulfone
 - (J) Polyurethane and urea resins
 - (K) Polyamides and copolyamides derived from diamines and dicarboxylic acids and/or aminocarboxylic acids or Nylon 6, nylon 66, nylon 11, and nylon 12
 - (L) Polyamide-polyethers
- (M) Polyesters derived from dicarboxylic acids and dialcohols and/or hydroxycarboxylic acids or the corresponding 50

Polyethylene terephthalate, polybutylene terephthalate and poly(1,4-dimethylolcyclohexane terephthalate)

- (N) Crosslinked polymers derived from aldehydes and phenois, urea or melamine Phenol/formaldehyde resin, urea/formaldehyde resin, and melamine/formaldehyde resin
- (O) Alkyd resins Glycerin/phthalic acid resin.

(P) Unsaturated polyester resins derived from copolyesters of saturated and unsaturated dicarboxylic acids and polyhydric alcohols using vinyl compounds as crosslinking agents, and halogen-containing modified resins thereof

(O) Natural polymers

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Cellulose, rubber and protein, or derivatives thereof, such as cellulose acetate, cellulose propionate, cellulose acetate and cellulose ether

When the copolymers of this invention are used as synthetic waxes, they may, of course, be mixed with various known waxes

The random copolymers of this invention may be used as a blend with each other.

The random copolymer of this invention may be modified by grafting thereto a monomer selected from the group consisting of (a) alpha, beta-unsaturated carboxylic acids and/or their derivatives. (b) styrenes. (c) organic silicon compounds having an olefinic unsaturated bond and a hydrolyzable group and (d) unsaturated epoxy compounds. The modified random copolymer obtained has the same excellent properties as the random copolymer of the invention and also especially good adhesion to metals and synthetic resins and good compatibility with other resins. The amount of the grafting monomer in the modified random copolymer is 0.1 to 50 parts by weight, preferably 0.5 to 35% by weight per 100 pans by weight of the random copolymer of the invention. The modified random copolymer usually has an intrinsic viscosity [η] of 0.03 to 20 dl/g, preferably 0.05 to 5 dl/g. Examples of the grafting monomer are:

Examples of the unsaturared carboxylic acids and their derivatives (a) include unsaturated carboxylic acids such as acrylic acid, methacrylic acid, alpha-ethylacrylic acid, maleic acid, fumaric acid, itaconic acid, citraconic acid, teirahydrophthalic acid, methyltetrahydrophthalic acid, endocis-bicyclo[2,2,1]-hept-5-ene-2,3-dicarboxylic acid (Nadic acid®), and methylendocis-bicyclo[2.2.1]-hept-5-ene-2,3-dicarboxylic acid (methylNadic acid®), and their nalices amides, imides, anhydrides and esters. Specific examples of these derivatives are malenyl chloride, maleimide, maleius annydride, citrasonic anhydride, monomethyl maleate and dimethyl maleate. Of these, the unsaturated dicardoxylis acids or their annydrides are preferred. Maleic acid. Nadic acid or their annydrides are especially preferred

Examples of the styrenes (b) are compounds represented by the following formula

$$R^{20} = C = R^{21}$$

wherein H²⁰, H²¹ and H²² each represent a hydrogen atom or a lower alkyl group. Specific examples are styrene, aicha-methylstyrene, o-methylstyrene, m-methylstyrene, p-methylstyrene, m-ethylstyrene, p-ethylstyrene, o-iso-propylsytrene, m-isopropylstyrene and p-isopropylstyrene. Of these, styrene, m-methylstyrene and p-methylstyrene are 40

Examples of the organic silicon compound (c) having an clefinic unsaturated bond and a hydrolyzable group are compounds represented by the following formula

wherein R23 and R24 are identical or different and each represents a monovalent group composed of carbon hydrogen and optionally oxygen and having an olefinically unsaturated bond, and Y^1 and Y^2 are identical or different ÷5 and each represents a hydrolyzable group. compounds of the following formula

wherein \mathbb{R}^{23} is as defined, and \mathbb{Y}^1 , \mathbb{Y}^2 and \mathbb{Y}^3 are identical and different and each represents a hydrolyzable group. Ξε and compounds of the following formula

wherein \mathbb{R}^{23} Y1 and Y2 are as defined above, and X represents an organic group free from an olefinic unsaturated

Examples of \mathbb{R}^{23} and \mathbb{R}^{24} are vinyl allyl, butenyl, cyclohexenyl, cyclopentadienyl, $\mathrm{CH_2=C(CH_3)COOO(CH_2)_{5^{-1}}}$ $\mathsf{CH_2} = \mathsf{C}(\mathsf{CH_3}) \mathsf{COO}(\mathsf{CH_2})_2 - \mathsf{O} - (\mathsf{CH_2})_3 - \text{ and } \mathsf{CH_2} = \mathsf{C}(\mathsf{CH_3}) \mathsf{COOCH_2OCH_2OCH_2OCH_2O(CH_2)_3} - \mathsf{Of these, the hydrocarbonic of the coordinate of the$

bon groups having a terminal olefinically unsaturated bond and esters thereof are preferred. The vinyl group is espe-

Examples of Y¹, Y² and Y³ are alkoxy groups such as methoxy, ethoxy, butoxy and methoxyethoxy, acyloxy groups such as formyloxy, acetoxy and propionoxy, oxime groups such as -ON=C(CH₃)₂, -ON=CHCH₂C₂H₅ and -ON=C $(C_6H_5)_2$, and substituted amino groups and arylamino groups such as -NHCH₃, -NHC₂H₃ and -NH(C₆H₃).

Examples of X in the above formula include methyl, ethyl, propyl, tetradecyl, octadecyl, phenyl, benzyl and toly: Of these, monovalent hydrocarbon groups are preferred.

Organic silicon compounds preferably used in this invention are those of the second formula given above, particularly those in which the groups Y¹, Y² and Y³ are identical. Especially preferred are vinyltrialkoxysilanes such as vinyltrimethoxysilane, vinyltrietnoxysilane, vinyltris(methoxyethoxy)silane, vinylmethyldiethoxysilane and vinylphe-

The unsaturated epoxy monomers (d) are preferably monomers having at least one polymerizable unsaturated bond and at least one epoxy group per molecule. Examples include unsaturated glycidyl esters represented by the

wherein R²⁵ represents a hydrocarbon group having a polymerizable ethylenically unsaturated bond.

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unsaturated glycicyl etners represented by the general formula

wherein R^{25} is as defined above, and Z represents a divalent group of the formula -CH₂-C- or

and epoxyalkenes represented by he general formula

wherein R^{25} is as defined, and R^{26} represents a hydrogen atom or a methyl group

Specific examples of the monomers (d) are glycicyl acrylate, giyoidyl methacrylate, mono- and di-glycidyl itaconate mono-, di- and-triglycidy! butene-tricarboxylates, mono- and di-glycidy! citraconates, mono-and di-glycidy! endo-cisbicyclo[2.2.1]-nept-5-ene-2.3-carboxylates, mono- and di-glycloyl endo-cis-bicyclo-[2.2.1]-hept-5-ene-2-methyl-2.3-dicarboxylates, mono-and di-glycidyl allylsuccinates, giycidyl p-styrene-carboxylate, allyl glycidyl ether, 2-metnylalkyl glycidyl ether, styrene p-glycidyl ether, 3,4-epoxy-1-butene, 3,4-epoxy-3-methyl-1-butene, 3,4-epoxy-1-pentene 3.4-3.4-epoxy-3-methyl-1-pentene, 5,6-epoxy-1-hexene, and vinyloydionexene monoxide. Of these grydidyl adrylate

The modified random copolymer may be produced by various known methods. For example, the random copolymer is melted, and the grafting monomer is added and graft-copolymerized. Alternatively, the random copolymer is dissolved in a solvent, and the grafting monomer is added and graft-copolymerized. In either case, the reaction is preferably carried out in the presence of a radical initiator in order to perform graft copolymerization efficiently. The radical initiator may include organic peroxides, organic peresters and azo compounds. Ionizing radiations and ultraviolet light may

The modified random copolymer obtained by the above procedure has excellent heat resistance as snown by its high glass transition temperature which is one measure of heat resistance. Specifically, the modified random copolymer usually has a glass transition temperature, measured by a Dynamic Mechanical Analyzer (DMA), of 20 to 250°C, in many cases 30 to 220°C. The modified random copolymer also usually has a softening temperature of 20 to 230°C. in many cases 30 to 200°C. The softening termperature is defined as the temperature at which a quartz needle (diameter

0.635 mm) penetrates 0.1 mm into the copolymer under a load of 49 g when the tamperature is elevated at a rate of 5°C/min. using TMS (thermomechanical analyzer, a product of E. I. du Pont de Nemours & Co.).

The modified copolymer usually has a density, determined by the method of ASTM D 1505, of 0.95 to 1.20 g/cm³ in many cases 0.95 to 1.10 g/cm³.

When the random copolymer of this invention is crosslinked with sulfur or organic peroxides or by actinic light such as electron beams or radioactive rays, its heat resistance, chemical resistance, solvent resistance and mechanica.

The properties of the cycloolefin random copolymers obtained in this invention were measured in accordance with the following method

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(1) Copolymer composition (mole%)

On the basis of the cycloolefin unit content of the random copolymer determined by 13°-NMR (200 MHz), a calibration curve is prepared between the cycloplefin unit content and the height of the peak of an absorption band based on the cycloolefin unit determined by infrared spectroscopy. The peak height of the absorption band of the infrared absorption spectrum of the random copolymer is measured, and the cycloolefin unit content is calculated The ethylene unit content of the random copolymer is calculated by subtracting the cycloolefin unit content from (2) Intrinsic viscosity [η]

Measured at 135°C using a Ubbelonde viscometer.

(3) Glass transition temperature (Tg)

Measured by a Dynamic Mechanical Analyzer (DMA, made by E. I. du Pont de Nemours & Co.).

(4) Crystallinity

Determined by X-ray diffractometry.

(5) Voiatile component (VM)

A weight change was measured under the conditions of 300°C, 1 torr and 1 hour.

(6) Content of the unreacted cycloolefin

The polymer was dissolved in cyclchexane, and the amount of the unreacted cyclcolefin was measured by gas chromatography (7) Molecular weight distribution

The molecular weight distribution ($\overline{\mathbb{M}}$ w/ $\overline{\mathbb{M}}$ n) is determined in the following manner in accordance with the procedure described at pages 14-114 of "Get Permeation chromatography" by Takeuchi, published on May 20, 1976

(1) Using standard polystyrene of a known molecular-weight (mono-dispersed polystyrene, a product of Toyo Soda 35 Mfg. Co., Ltd., Japan), the molecular weight M of the polystyrene sample and its GPC (gel permeation chromatography) count are measured. A calibration curve for the molecular weight M and the EV (elution volume) is drawn The concentration of the polymer at this time is set at 0.02% by weight. (2) A gel permeation chromatogram of the sample is taken by the GPC measuring method, and the number average 4.7

molecular weight (Mn = $\frac{\Sigma MN_L}{\Sigma N_L}$) and the weight average molecular weight (Mw = $\frac{\Sigma ML^2N_L}{\Sigma MN_L}$) are calculated for copolymer

by (1) above, and the value $(\overline{M}w/\overline{M}n)$ is determined

The sample is prepared under the following conditions, and the conditions for GPC are as shown below

45 Dreparation of the sample

- (a) The sample is taken into an Erlenmeyer flask together with o-dichlorobenzene so as to provide a 0 02% solution
- (b) into the Erienmeyer flask containing the sample is added 0.1% by weight, based on the polymer solution, of
- (c) The Erlenmeyer is heated to 140°C, and the contents are stirred for about 30 minutes to dissolve the polymer
- (d) Then, at 135 to 140°C, the solution is filtered by a 1η Millipoere filter.
- (e) The filtrate is subjected to gel permeation chromatography

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Conditions for get permeation chromatography

(a) Device. Model 150C, made by Waters Company

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- (b) Column: TSK-GMH6 (mix type) made by Toyo Soda Mfg. Co., Ltd.
- (c) Amount of the sample: 500 μl
- (d) Temperature: 140°C
- (e) Flow rate: 1 ml/min. (f) Total theoretical stages of the column;
- 1×10^4 to 4×10^4 (measured with acetone)

The following Examples further illustrate the present invention.

10 EXAMPLE 1

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Ethylene and 1.4.5.8-dimethano-1,2.3.4.4a.5.8.6a-octahydronaphthalene (Table 1, (a), appreviated as DMON] were copolymerized in a 2-liter polymerization reactor equipped with agitating blades.

The polymerization reactor was continuously charged at its top with a cyclonexane solution of DMON at a rate of 0.4 liter/hour so as to provide a DMON concentration of 60 g/liter in the polymerization reactor, a cyclohexane solution of VO(OC₂H₅)Ci₂ as a catalyst at a rate of 0.7 liter/hour so as to provide a vanadium concentration of 0.7 mmole/liter in the polymerization reactor (the concentration of variadium fed at this time was 2.86 times that in the polymerization reactor), a cyclchexane solution of ethyl aluminum sesquichioride as a catalyst at a rate of 0.4 liters/hour so as to provide an aluminum concentration of 5.6 mmoles/liter in the polymerization reactor, and 0.5 liters/hr of cyclonexane In the mean-time, the polymerization mixture in the polymerization reactor was withdrawn continuously from the bottom of the polymerization reactor so that the amount of the polymerization mixture in the reactor was always kept at 1 liter Furthermore, 80 liters/hr of ethylene, 80 liters/hr of nitrogen and 0.2 liter/hr of hydrogen were fed from the top of the polymerization reactor. The copolymerization was carried out at 10°C by circulating a cooling medium into a jacket fitted to the exterior portion of the reactor. As a result of the copolymerization reaction under the above conditions, a polymerization reaction mixture containing the ethylene/DMON random copolymer was obtained. A mixture of cyclonexane and isopropanol (1:1) was added to the polymer solution with drawn from the bottom of the polymerization reactor to stop the polymerization reactor. Thereafter, an aqueous solution composed of 1 liter of water and 5 ml of concentrated hydrochloric acid and the polymer solution were contacted at a ratio of 1:1 with strong stirring by a homomixer to transfer the catalyst residue to the aqueous layer. The mixture was left to stand, and the water layer was removed The residue was washed twice with distilled water to purify and separate the polymer solution. The polymer solution was contacted with 3 times its amount of acetone with strong stirring, and the solid ponton was collected by filtration and fully washed with acetone. Thereafter, the resulting solid portion was put into acetone in a concentration of 40 gr liter, and reacted at 60°C for 2 hours. The solid portion was then collected by filtration, and cried in a stream of nitrogen

By the above procedure, the ethylene/DMON copolymer was obtained at a rate of 94 g/hr. The copolymer had an ethylene content of 61.3 mole%, an intrinsic viscosity of 0.85, an Mw/Mn, determined by GPC, of 2.50, a crystallinity determined by X-ray diffractometry, of 0%, and a glass transition temperature (Tg) of 143°C. The copolymer contained 0.4% by weight of volatile compnents and 0.13% by weight of the unreacted monomers

EXAMPLES 2-14

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Example 1 was repeated except that the copolymerization conditions indicated in Table 2 were used. The properties of the resulting copolymers are shown in Table 3.

			T					
5			-0	octa-	1,8a-		141-	
10			1,4,5,8-dimethano-1,2,3,4,4a,5,8,8a-octahydro-	2-ethyl-1,4,5,8-dimethano-1,2,3,4,4a,5,8,8a-octa-	2,3-dimethyl-1,4,5,8-dimethano-1,2,3,4,4a,5,8,8a-octahydronaphthalene	2-methyl-1,4,5,8-dimethano-1,2,3,4,4a,5,8,8a- octahydronaphthalene	12-methylhexacyclo[6,6,1,1 ³ ,6,1 ^{10,13} ,0 ^{2,7} ,0 ^{9,14}]-	
:5			4a,5,8,8a	1,2,3,4,4	Jano-1,2,	1,2,3,4,4	6,110,13	
20	~ 1	lefin	-1,2,3,4,	methano-	,8-dimeth	imethano-	6,6,1,1 ³	
25	Table 1	Cycloolefin	imethano- ne	,4,5,8-di thalene	nyl-1,4,5 naphthale	,4,5,8-d; aphthaler	exacyclo[
30			l,4,5,8-dim naphthalene	2-ethyl-1 1ydronaph	2,3-dimethyl-1,4,5,8 octahydronaphthalene	2-methyl-l,4,5,8-dim octahydronaphthalene	12-methylhexa 4-heptadecene	
35				2115		3	CII3 12	
40					CH ₃	CII		
			(a)	(b)	(c)	(p)	(c)	l ,

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			1	i						<u> </u>						
5			-	ization tempera- turg (°C)		10	I	# :	:		= 1	<u>.</u> 20	i z	30	10	:
10			Amount	of hy- drogen fed (t/hr)		0.5	:	: :	2 2	ŧ	30	4 0.7	10	150	0.5	:
;5			Anount	othylene fed (t/hr)	S	9	•	: :	90) <u>.</u>	90	110	E	150	08 :	
20			punoc	Concentration in the reactor	(nmo]/e)	2.	£	= :	2 g	2.8	5.6	5.6		t 1		1
25	7	nditions	Altminum compound	Name	AI (C,IL), CI	1.5-1.5	x =		= =	=	= x	F :			A1 (C2115) 15 C1 1.5/	3) 2
30	Table 2	ation con			AI (C, II,	7								: :	A1 (C ₂ 11 ₅) A1 (C ₂ 11 ₅)	(7/3)
35		Copolymerization conditions	mpound	Concentration in the reactor	0.7	;	= r	ε ;		: :	: = 1	· _			:	
40		CoJ	Vanadium compound	Name	VO(0C2115)C12	:	: :	÷ =	= r	=	r r		r	אסכוי	VO(0C ₂ H ₅)C1,	
45			efin ^{*3)}	Concentration in the reactor (g/t)	09		r	: :	: :		• 00	2	E	09	>	
5 0			Cycloolefin	Name	Table 1	£ £	(O)	(g) :	(e) (a)		2 2	*		r	2	
is		Ex-			1	2	т.	4 10	9	g*1)	01	111	12 27	13	14	

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*1): The amount of nitrogen fed was 50 liters/hr. *2): Nitrogen was not fed. *3): Table 4

							:								
	Amount of the residual monomer (wt%)	0 13	0.12	0.12	0.12	0.13	0.21	0.12	0.11	0.15	0.11	0.14	0.11	0.10	0.12
	VM (Wt8)	0.4	0.5	0.4	0.5	0.5	9.0	0.5	0.3	9.0	0.5	0.5	0.2	0.3	0.4
polymer	Mw/An	2.5	2.3	2.6	2.4	2.7	2.9	2.3	3.0	2.7	2.5	2.4	3.0	2.9	2.4
the	DMβ-Т9 (°С)	143	135	140	147	127	155	152	131	137	9/	6/	45	160	145
	rystal- inity (%)	0	0	0	0 (0 (0 0	o (- ·	0	 0	0
Pro	[e]	0.85	0.82	0.83	48.0		/ () (, ,	15.0	2 2 2		0 0	0/•/	T/:	/9.
	Ethylene content (mol%)	61.3	64.8	2.50	0.4.0	4 4		• ~			6	`		· ·	0.00
	Amount of the copolymer (g/hr)			102	122	84	78	120	06	102	100	130	96	106	
Example		، ت	7 E	4	S	9	7	8 * 1)	6	10	11	12*2)	13	14	
	Properties of	Amount Ethylene [n] Crystal- DM8-Tg $\bar{M}\omega/\bar{M}n$ VM copolymer (mol%) (%)	Amount Ethylene [n] Crystal- $0M8^{-T}g$ $\overline{Mw/\overline{M}}n$ VM copolymer (mol%) [%) (%)	Amount Ethylene [n] Crystal- DM8-Tg Mw/M n VM copolymer (mol%) [%) [%) [%] [%] [%] [%] [%] [%] [%] [%] [%] [%]	Amount Ethylene (n) Crystal- DM8-Tg Mw/Mn VM (of the copolymer (mol%) (%) (%) (%) (%) (%) (%) (%) (%) (%) (Amount Ethylene (n) Crystal- DM8-Tg Mw/Mn VM (of the content (mol%) (%) (%) (%) (%) (%) (%) (%) (%) (%) (Amount Ethylene (n) Crystal- DM8-Tg Mw/Mn VM (of the content (mol%) (%) (%) (%) (%) (%) (%) (%) (%) (%) (Amount Ethylene (n) Crystal- (%C) MW-TG (wt%) (w	Amount Ethylene [n] Crystal- DM8-Tg Mw/Mn Amount of copolymer (mol%) 104 61.3 0.85 0 143 2.5 0.4 0.12 105 64.8 0.82 0 140 2.6 0.4 107 64.8 0.84 0 140 2.6 0.4 108 65.6 0.91 0 147 2.4 0.5 108 65.6 0.91 0 155 2.9 0.6 120 67.0 0.89 0 155 2.9 0.6 121 2.7 0.5 0.13	Amount Ethylenc [n] Crystal- DM8-Tg Mw/Mn VM Amount of copolymer (mol%) [%] [%] [%] [%] [%] [%] [%] [%] [%] [%]	Amount of the copolymer of the copolymer of the copolymer (mol%) Ethylene content (mol%) (mo	Amount of the content of full copolymer Ethylene content (mol8) (mol8) Crystal linity (g/hr) DM8-Tg (wt8) Mw/Hn (wt8) Amount of the casidual monomer (wt8) 104 61.3 0.85 0 143 2.5 0.4 0.13 96 64.8 0.85 0 143 2.5 0.4 0.13 102 64.8 0.85 0 145 2.6 0.4 0.13 102 64.8 0.83 0 140 2.6 0.4 0.12 102 64.9 0.84 0 147 2.4 0.5 0.12 102 64.0 0.84 0 147 2.4 0.5 0.12 122 70.2 0.91 0 147 2.4 0.5 0.13 84 67.6 0.57 0 0 152 2.9 0.6 0.13 120 67.2 0.31 0 131 3.0 0.5 0.11 102 <td< td=""><td>Amount of the continue content of the content of the content of the content (mols) Crystal of the copolymer (mols) Crystal of the copolymer (mols) Thinity (%C) MW/Mn VM Amount of the residual the residual monomer (Wts) 104 61.3 0.85 0 143 2.5 0.4 0.13 96 64.8 0.82 0 135 2.3 0.5 0.13 102 64.0 0.83 0 140 2.6 0.4 0.12 102 64.0 0.83 0 140 2.6 0.4 0.12 102 64.0 0.84 0 140 2.6 0.4 0.12 102 64.0 0.84 0 147 2.4 0.5 0.12 12 70.2 0.91 0 147 2.7 0.5 0.13 12 67.6 0.57 0 0 152 2.3 0.5 0.13 102 60.3 0.70 0 137 2.7 0.6 <td< td=""><td>Amount Ethylenc [1,1] Crystal- (R_0) (R_0)</td><td>Amount of the copolymer of the copolymer of the copolymer of the copolymer (mol%) (n) Crystal (%C) DHR-T9 (Wt%) Mw/Mm (Wt%) Amount of the residual (Wt%) copolymer (gd/hc) (mol%) (1101 ft) (1101 ft) (Mt%) (M</td></td<></td></td<>	Amount of the continue content of the content of the content of the content (mols) Crystal of the copolymer (mols) Crystal of the copolymer (mols) Thinity (%C) MW/Mn VM Amount of the residual the residual monomer (Wts) 104 61.3 0.85 0 143 2.5 0.4 0.13 96 64.8 0.82 0 135 2.3 0.5 0.13 102 64.0 0.83 0 140 2.6 0.4 0.12 102 64.0 0.83 0 140 2.6 0.4 0.12 102 64.0 0.84 0 140 2.6 0.4 0.12 102 64.0 0.84 0 147 2.4 0.5 0.12 12 70.2 0.91 0 147 2.7 0.5 0.13 12 67.6 0.57 0 0 152 2.3 0.5 0.13 102 60.3 0.70 0 137 2.7 0.6 <td< td=""><td>Amount Ethylenc [1,1] Crystal- (R_0) (R_0)</td><td>Amount of the copolymer of the copolymer of the copolymer of the copolymer (mol%) (n) Crystal (%C) DHR-T9 (Wt%) Mw/Mm (Wt%) Amount of the residual (Wt%) copolymer (gd/hc) (mol%) (1101 ft) (1101 ft) (Mt%) (M</td></td<>	Amount Ethylenc [1,1] Crystal- (R_0)	Amount of the copolymer of the copolymer of the copolymer of the copolymer (mol%) (n) Crystal (%C) DHR-T9 (Wt%) Mw/Mm (Wt%) Amount of the residual (Wt%) copolymer (gd/hc) (mol%) (1101 ft) (1101 ft) (Mt%) (M

EXAMPLE 15

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Using a 2-liter glass polymerization reactor equipped with agitating blades, ethylene and 2-methyl-1,4,5,8-dimeth-ano-1,2,3,4,4a,5,8,8a-octahydronaphtahlene (to be abbreviated as M-DMON, (d) of Table 1 were copolymerized continuously

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Specifically, the reactor was charged at its top with 0.9 liter/hr of a toluene solution of M-DMON so as to provide an M-DMON concentration in the reactor of 60 g/liter, 0.7 liter/hr of a toluene solution of VO(OC₂H₅)Cl₂ as a catalyst so as to provide a vanadium concentration in the reactor of 0.5 mmole/liter, and 0.4 liter/hr of a toluene solution of ethyl aluminum sesquichloride as a catalyst so as to provide an aluminum concentration in the reactor of 2 mmoles/liter in the meantime, the polymer solution was continuously withorawn from the bottom of the reactor so that the amount of the polymer solution in the reactor was maintained always at 1 liter. Furthermore, 35 liters/hr of ethylene and 60 liters, a cooling medium through a jacket fitted to the outside of the reaction was carried out at 10°C by circulating under the aforesaid conditions, a polymerization reaction mixture containing an ethylene/M-DMON random copolymerization to stop the copolymerization reaction. The polymer solution was then put into a large amount of acetone/methanol to precipitate the resulting copolymer. The copolymer was fully washed with acetone and dried under reduced pressure for one day at 80°C. By the above operation, the ethylen/M-DMON random copolymer was obtained at a rate of 60 g

The copolymer was found to have an ethylene content of 63 mole%. The copolymer had an intrinsic viscosity measured in decalin at 135°C, of 0.92, an iodine number of 1.0, a crystallinity, determined by X-ray diffractometry of 6, and a transparency, determined by a haze meter on a 1 mm sheet sample in accordance with ASTM D 1003-52 of 3.1%

The glass transition temperature Tg of the copolymer, determined from the peak temperature of the loss modulus E" which was measured at a temperature elevation rate of 5°C/min, by a dynamic mechanizal analyzer was 127°C list melting point Tm was measured within a temperature range of -120 to 400°C by DSC (Du Pont 990 type) at a temperature elevation rate of 10°C/min.) No melting curve (peak) was observed

EXAMPLES 16-24 AND COMPARATIVE EXAMPLE 1

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Example 15 was repeated except that the copolymerization conditions shown in Tables 4 and 5 were used. The properties of the resulting copolymers are shown in Table 5. In Example 22, a 1/1 mixture of toluene and cyclonexane was used as the polymerization solvent.

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			Ī		T-											
5			nd	Concentration in the reactor (mmol/t)	2.0	£	2	6.4	2	4.0	6.4	1				: :
15			nodwoo u		C1 _{1.5}								-			
15			Aluminum compound	Name	A1 (C2115) 1.5 C1 1	=	ž	=	:	*	=	Ε	Ε	=	=	
23					+											
25	Table 4	Catalyst	punodwoa	Concentration in the reactor (mmol/k)	0.5	z	=	0.8	=	0.5	0.8	=	=	ī	8.0) -
30 -	Tab	20	Vanadium compound	Мате	VO(OC2H5)C12	Ł	r	=	voc1,	VO(OC, III,) C.1,	7	ŧ	2	=	=	
42		Cycloolefin		Concentration in the reactor (g/t)	09	E	:	E	=	9	09	ı	E	Ε	09	
48				Name	Table 1 (d)	z	*	E		=	(p)	(c)	(a)	(e)	(p)	
50		Example			15	16	17	18	19	20	21	22	23	24	Comp Ex. 1	

									_						
5		ñw∕Ñn		2.5	2.8	ה ה	2.0	٤٠٠	7.7	2.8	3.0	2.9	2.5	1	_
		185-1111 (2C)			 I i	l 1	1	02	2	ı	 I	1	1	1	
7.3	l yme r	(2°)	761	118	107	116	115	34	5 = =		777	115	131	1	_
15	ocloo a	Haze (8)	-	7.7	2.0	2.2	2.3	5.5	9 6	٠, د	٠٠, و	2.4	2.8	ı	_
co	Properties of the copolymer	Crystal- linity (%)	0	0	0	0	0	0	0	0	> <	 ->	0	1	
25	Proper	Iœline value	1.0	0.9	6.0	0.7	0.0	1.0	6.0	0.8	9 0	n ,	1.0	1	_
e Table 5		Ξ	0.92	1.21	1.40	1.25	1.23	2.94	1.18	1.20	- 10	1 1 0	رد. د	1	
30 E		Ethylene content (mole%)	63	99	72	89	1.9	94	67	99	65	. 09	0	ı	
35	Anount of the	copolymer yielded (v/hr)	09	94	140	106	92	64	90	98	92	7.0		trace	-
45															
45		turg (°C)	10	E	2	ε	=	30	10	=	r	=		10	
£2	Amount of	fed (t/ht)	35	09	100	09	=	90	09	<u> </u>	=	r		09	
	Ex- ample		15	16	17	18	19	50	21	22	23	24	CIMO.	× .	i

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EXAMPLE 25

Fifty grams of the ethylene/DMON random co-polymer (A) obtained in Example 1 was dissolved in 500 ml of xylene at 125°C in a nitrogen atmosphere in a glass reactor. Then, a xylene sollution of maleic anhydride (abbreviafted as MAH) (12 g/30 ml) and a xylene solution of dicumyl peroxide (abbreviated as DCP) (0.27 g/30 ml) were fed gradua... over a period of 4 hours from separate lines into the reactor

Since MAH had low solubilty in xylene, it was dissolved by heating with an infrared lamp, and the heating was continued until its feeding was terminated

After the feeding, the reaction was continued for 2 hours, and after the reaction, the reaction solution was cooled to room temperature. The reaction solution was put into a large amount of acetone to precipitate the polymer. The resulting grafted ethylene/DMON copolymer was filtered, repeatedly washed with acetone, and dried under reduced pressure at 50°C for 1 day to give the desired MAH-grafted ethylene/DMON copolymer. The amount of MAH grafted measured by oxygen analysis of the modified copolymer was 1.13 by weight. The modified copolymer had an intrins of viscosity, measured in decalin at 135°C, of 0.85 dl/g, a density, determined in accordance with ASTM D 1505, of 1.025 groms, a grass transition temperature (Tg) of 145°C

EXAMPLES 25 & 27

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Example 25 was repeated except that 50 g of each of the cycloclefin random copolymers indicated in Table 5 was used instead of the ethylene/DMON random copolymer, and the grafting conditions were changed as in Table 6. The

 'n	10	_
 _	ıe	-

Grafting (AH (g/30 ml)	COR (e/30 mix
AH (g/30 ml)	DOP (5/30 mi)
	1 20. (9.30 (11)
12	0.27
12	0 27
	12

Table 7

Example	Properties	s of the	modified copolymer	
	Amount of MAm grafted (wt.%)		1 - 0 -	Density (gram ³)
26	1 13	0.88	145	1.025
27	1.23	0.86	140	1 023

EXAMPLES 28-30

Example 25 was repeated except that the random copolymer, grafting monomer and the radical initiator were changed as shown in Table 8. The results are shown in Table 9.

Table 8

	Random occosyme:	Amount (g)	Gratting monomer	Amount (a)	DOP (c)
25	<u>l</u> A	50	a	20	0.33
29	D	50	ε	16	0.34
30 ———	ΕΕ	50	ε	10	0.27

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Table 6 (continued)

Example	Random copolymer	Amount (g)	Grafting monomer		
A: Obt	ained in Example 1.		- and griffortonier	Amount (g)	DCP (g
	stained in Example 2, a	nd			
	tained in Example 3				
	yl-tri-ethoxy silane				
b: sty					
C: all	cidyl methacrylate				

PalcsT

Example	Properties o	of the mo	odified copolymer	
	Amount of grafted monomer (wt.%)	[η]	Tg (by DMA method)	Density (g/cm³)
25	0.25	0.87	144	1.026
29	1 34	0.86	:38	
30	1 13	0.85	142	1.021

Claims

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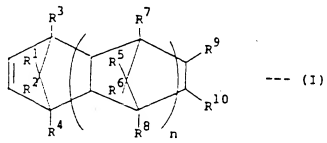
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1. A random copolymer which is transparent and is characterized in that

(A) the copolymer comprises polymerized units of ethylene and polymerized units of at least one cycloolefin represented by the following formula (I)



wherein \mathbb{R}^1 , \mathbb{R}^2 , \mathbb{R}^3 , \mathbb{R}^4 , \mathbb{R}^5 , \mathbb{R}^7 and \mathbb{R}^5 are identical or different and each represents a hydrogen atom has open atom or a hydrogen group having 1 to 10 carbon atoms, and \mathbb{R}^9 and \mathbb{R}^{10} are identical or different or \mathbb{R}^9 and \mathbb{R}^{10} are identical or different or \mathbb{R}^9 and \mathbb{R}^{10} are bonded to each other to form a group of the following formula

$$(R^{11}-C-R^{12})_{\mathfrak{g}}$$

in which R¹¹ and R¹² are identical or different and each represents a hydrogen atom, a halogen atom or a hydroparbon group having 1 to 10 carbon atoms, and *E* is an integer of 3 or 4, and n is a positive integer of 1 to 3 (C) the mole ratio of the polymerized cycloplefin units to the polymerized ethylene units is from 3:97 to 60:40, by the creach cycloplefin of formula (I) is incorporated in the polymer chain as polymerized units represented by the following formula (II)

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wherein all symbols are as defined above.

- (D) the copolymer has an intrinsic viscosity [n], measured in decalin at 135°C, of 0.01 to 20 dl/g.
- (E) the copolymer has a molecular weight distribution (Mw/Mn), measured by gel permeation chromatography, of not more than 4.
- (F) the copolymer has a crystallinity, measured by X-ray diffractometry, of 0 to 10%, and
- (G) the copolymer has a glass transition temperature (Tg) of 10 to 240°C as measured by a Dynamic Mechanical Analyzer.
- 2. A random copolymer according to claim 1 wherein the mole ratio (B) of the polymerized cycloolefin units to the polymerized ethylene units is from 5:95 to 50:50.
 - 3. A random copolymer according to claim 1 or 2 which has a glass transition temperature (Tg) of 20 to 200°C
- A process for producing a random copolymer as defined in any one of the preceding claims, which process comprises continuously copolymerizing ethylene with the or each cycloolefin represented by formula (i) in a hydrocarbon medium in the presence of a catalyst formed from a vanadium compound soluble in the hydrocarbon medium and an organo-aluminum compound while maintaining the atomic ratio of aluminum atoms to vanadium atoms in the reaction system, A1:V, at 2 or more:1.

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- 5. A process according to claim 4 wherein the soluble variadium compound is added to the reaction system in a concentration not more than 10 times its existing concentration in the reaction system in order to maintain the atomic ratio At:V at 2 or more:1.
- 6. A process for preparing a modified random copolymer, which process comprises grafting on to a random copolymer, as defined in any one of claims 1 to 4, 0.1 to 50 parts by weight of a monomer selected from (a) alona, beta-unsaturated carboxylic acids and/or their derivatives. (b) styrenes, (c) organic scioon compounds having an olefinically unsaturated bond and a hydrolyzable group, and (d) unsaturated epoxy monomers per 100 parts by weight of the said random copolymer.

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- 7. Molded anicles prepared from a random copolymer as defined in any one of claims 1 to 4 or from a modified random copolymer prepared by a process as claimed in claim 6.
- 8. A modified random copolymer which is obtainable by a process as defined in claim 6.

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Patentansprüche

1. Ein statistisches Copolymer, das transparent ist und dadurch gekennzeichnet ist, daß

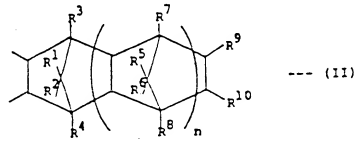
(A) das Copplymer polymerisierte Einneiten von Ethylen und polymerisierte Einheiten von wenigstens einem Cycloblefin umfaßt, das durch die folgende Formel (I) dargestellt ist

worin R¹, R², R³, R⁴, R⁵, R⁶, R⁷ und R⁵ gleich oder verschieden voneinander sind und jeweils ein Wassers, stoffatom, Halogenatom oder eine Kohlenwasserstoffgruppe mit 1 bis 10 Kohlenstoffatomen darstellt und R⁵ und R¹⁰ gleich oder verschieden voneinander sind und jeweils ein Wasserstoffatom, ein Halogenatom oder eine Kohlenwasserstoffgruppe mit 1 bis 10 Kohlenstoffatomen darstellen oder R⁹ und R¹⁰ miteinander verbunden sind, um eine Gruppe der folgenden Formel

$$(R^{11} - C - R^{12})_{t}$$

zu bilden, worin \mathbb{R}^{11} und \mathbb{R}^{12} gleich oder verschieden voneinander sind und jeweils ein Wasserstoffatom, ein Halogenatom oder eine Kohlenwasserstoffgruppe mit 1 bis 10 Kohlenstoffatomen darsteilen und 1 eine ganze Zahl von 3 bis 4 ist und n einen positive ganze Zahl von 1 bis 3 ist;

- (B) das Molverhältnis der polymerisierten Cycloolefineinneiten zu den polymerisierten Ethyleneinneiten von 3.97 bis 50:40 betragt;
- (C) das oder jedes Cycloolefin der Forme! (I) in die Polymerkette als polymerisierte Einneiten eingebracht sind, welche durch die folgende Forme! (II) dargestellt sind



worm alle Symbole die open angegebene Bedeutung besitzen

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- (D) cas Copolymer eine in Decatin bei 135°C gemessene intrinsiche Viskosität (η) von einschließlich 0.01 bis 20 d $^{\prime}$ g besitzt
- (E) das Copolymer eine durch Gelpermeationschromatographie gemessene Molekulargewichtsveriellung (Mw/Mh) von nicht mehr als 4 besitzt.
- (F) das Copolymer eine aurch Röntgenbeugung gemessene Kristallinität von einschließlich 0 bis 10 % besitzt und
- (G) das Copolymer eine mit einem dynamischen mechanischen Analysator gemessene Glasübergangstemperatur (Tg) von 10 bis 240°C besitzt.
- Statistisches Copolymer nach Anspruch 1, wobei das Molverhaltnis (B) der polymerisierten Cycloolefineinheiten zu den polymerisierten Ethyleneinheiten zwischen einschließlich 5,95 bis 50 50 liegt.
 - 3. Statistisches Copolymer nach Anspruch 1 oder 2, das eine Glasübergangstemperatur (Tg) von 20 bis 200°C auf-

weist.

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- 4. Ein Verlahren zur Herstellung eines wie in jedem der vorhergehenden Ansprüche definierten statistischen Copolymeres, wobei das Verlahren das kontinuierliche Copolymerisieren von Ethylen mit dem oder jedem Cycloolefin umfaßt, das durch die Formel (I) dargestellt ist, in einem Kohlenwasserstoffmedium in Gegenwart eines Katalysators, der aus einer in dem Kohlenwasserstoffmedium löslichen Vanadiumverbindung und einer Organo-Aluminiumverbindung gebildet ist, währenddessen das Atomverhältnis von Aluminiumatomen zu Vandiumatomen Al. Vin dem Beaktionssystem bei 2 oder mehr:1 gehalten wird.
- Verfahren nach Anspruch 4, worin die lösliche Vanadiumverbindung dem Reaktionssystem in einer Konzentration von nicht mehr als dem 10fachen seiner vorhandenen Konzentration in dem Reaktionssystem zugegeben wird um das Atomverhaltnis Al-V bei 2 oder mehr: 1 zu halten.
- 6. Verlahren zum Herstellen eines modifizierten statistischen Copolymeres, wobei das Verlahren, das Aufpfropfen auf ein wie in jedem der Ansprüche 1 bis 4 definiertes statistisches Copolymer von 0.1 bis 50 Gewichtstelle eines Monomeres pro 100 Gewichtstelle des statistischen Copolymeres umfaßt, das aus (a) α, β-ungesättigten Carbonsauren und/oder deren Dervlaten, (b) Styrolen, (c) organischen Siliconverbindungen mit einer olefinisch ungesättigten Bindung und einer hydrollisierbaren Gruppe und (d) ungesattigten Epoximonomeren ausgewahlt ist.
- Geformte Gegenstände, hergestellt aus einem wie in jedem der Ansprüche 1 bis 4 definierten statistischen Copolymer oder aus einem durch ein wie in Ansprüch 6 beansprüchtes Verfahren hergestelltes modifiziertes statistisches Copolymer
- 8. Ein modifiziertes, statistisches Copolymer, das durch ein wie in Anspruch 6 definiertes Verfahren erhältlich ist.

Revendications

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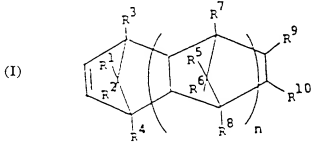
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- 1. Copolymère statistique transparent caractérisé en ce que
 - (A) le copolymère comporte des motifs polymérisés dérivés d'éthylène et des motifs polymérisés dérivés d'au moins une cyclopiéfine représentée par la formule suivante (I)



cans laquelle R1, R2, R3, R4, R5, R5, R7 et R5, identiques ou différents, représentent chadun un atome d'hydrogène, un atome d'halogène ou un groupe hydroparboné comportant de 1 à 10 atomes de carbone, et R5 pe hydrocarboné comportant de 1 à 10 atomes de carbone, et R5 pe hydrocarboné comportant de 1 à 10 atomes de carbone, ou R5 et R10 sont liés l'un à l'autre pour former un groupe de formule.

cans laquelle R¹¹ et R¹², identiques ou différents, représentent chacun un atome d'hydrogène, un atome d'haiogène ou un groupe hydrocarboné componant de 1 à 10 atomes de carbone, et 1 est un nombre entier égal à 3 ou 4, et n'est un nombre entier positif compris entre 1 et 3;

(B) le rapport molaire des motifs polymérisés dérivés d'une cyclooléfine aux motifs polymérisés dérivés d'éthy-

lène est compris entre 3/97 et 60/40;

(C) la cyclooléfine ou chaque cyclooléfine de formule (I) est incorporce dans la chaîne du polymère sous forme de motifs polymérises représentés par la formule suivante (II)

dans laquelle tous les symboles sont définis ci-dessus:

- (D) le copolymère à une viscosité intrinsèque [η], mesurée dans la Décaline® à 135 °C, comprise entre 0.00 et 20 dl/g;
- (E) le copolymère à un indice de polydispersité (M_p/M_n), mesuré par chromatographie par perméation de gel, ne dépassant pas 4;
- (F) le copolymère a un taux de cristallinité, mesuré par diffraction de rayons X, compris entre 0 et 10 %;
- (G) le copolymère a une température de transition vitreuse, déterminée avec un appareil d'analyse mécanicodynamique, comprise entre 10 et 240 °C.
- Copolymère statistique conforme à la revendication 1 dans lequel le rapport molaire (B) des motifs polymérisés dérivés d'une cyclooléfine aux motifs polymérisés dérivés d'éthylène est compris entre 5/95 et 50/50.
 - 3. Copolymère statistique conforme à la revendication 1 ou 2 ayant une température de transition vitreuse (T_v) comprise entre 20 et 200 °C.
- 4. Procédé de préparation d'un copolymère statistique conforme à une queiconque des revendications précédentes, lequel procédé comprend la copolymérisation en continu d'éthylène et de la ou de chaque cyclooléfine représentée par la formule (!), dans un milieu hydrocarboné en présence d'un catalyseur formé à partir d'un composé de vanadium soluble cans le milieu hydrocarboné et d'un composé organo-aluminique, avec maintient du rapport des atomes d'aluminium aux atomes de vanadium AI/V dans le système réactionne! à une valeur supérieure ou égale à 2/1.
 - Procédé conforme à la revendication 4 dans lequel on ajoute au système réactionnel le composé de vanacium soluble en une concentration n'excédant pas 10 fois sa concentration existant dans le système réactionnel afin de maintenir un rapport atomique AI/V supérieur ou égal à 2/1.
 - 6. Procédé de préparation d'un copolymère statistique modifié, lequel procédé comprend le greffage de 0,1 à 50 parties en poids, par rapport à 100 parties en poids dudit copolymère statistique, d'un monomère choisi parmi (a) les acides carboxyliques α,β-insaturés et/ou leurs dérivés, (b) les styrènes, (c) les composés organiques qu silicium comportant une liaison à insaturation éthylénique et un groupe hydrolysable, et (d) des monomères insaturés à fonction époxyde, sur un copolymère statistique défini par une quelconque des revendications 1 à 4.
 - Articles moulés préparés à partir d'un copolymère statistique conforme à une quelconque des revendications 1 à 4 ou à partir d'un copolymère statistique modifié préparé selon un procédé conforme à la revendication 6.
- 50 8. Copolymère statistique modifié pouvant être obtenu selon un procédé conforme à la revendication 6

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